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(54) Title: GENETICALLY MODIFIED CELLS EXPRESSING A TGF-BETA INHIBITOR, THE CELLS BEING LUNG CANCER CELLS

(57) Abstract: The present invention relates to compositions comprising a therapeutically effective amount of genetically modified cells containing a genetic construct expressing a  $TGF\beta$  inhibitor effective to reduce expression of  $TGF\beta$ , where the genetically modified cells are non-small cell lung cancer (NSCLC) cells or small cell lung cancer (SCLC) cells, and related methods.

# GENETICALLY MODIFIED CELLS EXPRESSING A TGF $\beta$ INHIBITOR, THE CELLS BEING LUNG CANCER CELLS Background of the Invention

#### Field of the Invention

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The present invention relates to compositions comprising a therapeutically effective amount of genetically modified cells containing a genetic construct expressing a  $TGF\beta$  inhibitor effective to reduce expression of  $TGF\beta$ , where the genetically modified cells are non-small cell lung cancer (NSCLC) or small cell lung cancer (SCLC) cells, and related methods.

#### Description of the Related Art

Lung cancer remains the most prevalent cancer in the western world, accounting for 30% of all cancerrelated deaths (Ramanathan and Belani, 1997). The current prognosis for patients with lung cancer is poor. The overall
cure rate is estimated as low as 13%. Approximately 180,000 new cases of lung cancer are expected in the United
States in 1999. The majority of these patients will die of their disease with 160,000 deaths from lung cancer
expected nation-wide in 1999.

There are two major subdivisions of lung cancer: 1) non-small cell (NSCLC) and 2) small cell lung cancer (SCLC). Treatment approaches and natural history differ for these two diseases. The majority (80%) of cases of lung cancer in the United States are NSCLC. Although advances in the understanding of important clinical and prognostic factors for both NSCLC and SCLC have been made in the past 20 years, there have been minimal improvements in therapeutic results. The only curative option for patients with NSCLC is local therapy (surgical excision or local irradiation) in patients with early stage disease (I & II) when the tumor is still localized. At diagnosis however, the majority of patients with NSCLC present with advanced disease, which is not curable by surgery alone. In advanced stages of disease, systemic chemotherapy and/or irradiation can produce objective responses and palliation of symptoms, however, they offer only modest improvements in survival. The median survival of patients with non-resectable disease is 6-12 months. Two-year survival rates for stages IIIB and IV NSCLC are 10.8 and 5.4 percent respectively. Likewise, five-year survival rates are 3.9 and 1.3 percent. Recently, several new drugs have become available for the treatment of NSCLC including paclitaxel (Taxol), docetaxel (Taxotere), topotecan, irinotecan, vinorelbine, and gemcitabine. While these drugs are improvements over prior chemotherapeutic agents (etoposide, cisplatin and carboplatin), the overall cure rate remains low.

SCLC is a very aggressive cancer which metastasizes early and often, and it has a median survival from diagnosis of only two to four months. Localized forms of treatment, such as surgical resection or radiation therapy, rarely produce long-term survival because of this cancer's propensity for distant metastasis. With chemotherapy, survival can be prolonged at least four to five times the media survival rate for patients who are given no therapy, however the overall survival at five years remains at only 5-10%.

Since current therapeutic modalities do not significantly enhance life expectancy in stages of NSCLC or SCLC patients, exploration of new therapeutic approaches for these patients is justified.

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#### Summary of the Invention

Patients bearing tumors of different histologic origin have elevated levels of Transforming Growth Factors- $\beta$ s (TGF $\beta$ s). TGF $\beta$ s are growth factors that are associated with immunosuppression. Suppression of the patients' immune system results in their inability to recognize and destroy tumors when they first appear. Furthermore, suppression of patients' immunity makes them susceptible to frequent infections. Injection of genetically engineered tumor cells to block their TGF $\beta$  production makes the gene modified cells potent vaccines that are recognized by and can activate the immune system against the tumor. Activation of the immune system subsequently causes the recognition and control of the parental, unmodified tumors in the host organisms. This phenomenon applies in animal tumor models and in human clinical trials. Thus, we propose to use this approach in patients with stages of non-small cell lung and small cell lung cancers.

# **Detailed Description of the Preferred Embodiment**

Lung cancers account for 30% of all death due to cancer in the United States (Ramanathan and Belani, 1997). The overall cure rate for lung cancer is 13% and the current prognosis for patients with non-small cell lung (NSCLC) and small cell lung cancer (SCLC) remains poor.

It has been documented that patients with progressive tumor growth have impaired immune function (Jakowlew et al. 1995, Ransohoff et al 1991; Holladay et al. 1992a; Holladay et al. 1992b). This impairment, commonly characterized as marked immune hyporesponsiveness, is not solely confined to tumor specific immunity, but rather, is often observed throughout the immune system. Impairment is especially evident in the cell-mediated or T-cell compartment and is characterized by T-cell lymphopenia and impaired T-cell responsiveness to both tumor specific and non-tumor specific stimuli (Ransohoff et al. 1991). One way tumors may escape immune surveillance is by expressing lower levels of MHC Class I and Class II molecules. Other tumors may escape by increasing the expression of immunosuppressor molecules, such as the TGFβs. It is common to observe tumors utilizing a combination of these mechanisms.

Gene therapy has received considerable attention in recent years. Vaccination with tumor cells designed to augment tumor antigen presentation and induce specific anti-tumor immunity has yielded promising but limited results (Holladay et al. 1992). Advances in our understanding of cancer biology and developments in vector technologies are advancing the therapeutic potential of tumor vaccine approaches. It is now possible to genetically modify tumor cells for vaccination to express specific tumor suppressor genes, immune modulators, drug sensitive genes and antisense gene fragments (Huber et al. 1991; Culver et al. 1992; Trojan et al. 1992; Dranoff et al. 1993; Ram et al. 1993; Trojan et al. 1993; Swisher et al. 1999). In particular preclinical and clinical studies demonstrate the potential of gene therapy approaches in treating lung cancer. Preclinical lung cancer models have shown regression of established tumors and enhanced immunogenicity using an allogeneic lung cancer line genetically modified to express the cytokine GM-CSF and a drug sensitive gene, herpes simplex virus thymidine kinase admixed with syngeneic bone-marrow derived dendritic cells (Miller et al. 1998). Preliminary results from phase I clinical trials in patients using retroviral gene therapy shows gene therapy to be well tolerated and without toxicity (Swisher et al. 1998).

Transforming growth factors beta (TGF $\beta$ ) are a family of multi-functional proteins that regulate the growth and function of many normal and neoplastic cell types (Sporn et al. 1986; Massague 1987; Border and Rouslahti 1992; Jachimczak et al. 1993). They exert a wide range of effects on a variety of cell types and have been shown to stimulate or inhibit cell growth, induce apoptosis and increase angiogenesis (Merzak et al. 1994; Jennings et al. 1994; Ashley et al.1998a; Ashley et al.1998b; Jennings et al. 1998). These effects are mediated at the level of signal transduction. TGF $\beta$  signal transduction has been found to affect the expression of more than 20 different genes (Baker and Harland 1997; Heldin et al. 1997; Stiles 1997; Yingling et al. 1997).

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TGF $\beta$  exists in three isoforms, known as TGF $\beta$ 1, TGF $\beta$ 2, and TGF $\beta$ 3. Their amino acid sequences display homologies on the order of 70-80%. Human TGF $\beta$  proteins and genes encoding them are known in the art. Specifically, TGF $\beta$ 1 mRNA (GenBank Accession No. XM\_008912 and NM\_000660), TGF $\beta$ 2 mRNA (GenBank Accession No. XM\_001754 and NM\_003238), and TGF $\beta$ 3 mRNA (GenBank Accession No. XM\_007417) from human sources have been documented.

TGF $\beta$  receptor proteins may be type I (55kDa) or type II (70kDa). TGF $\beta$  receptor proteins and genes encoding them are also known in the art. Human TGF $\beta$  receptor type I mRNA (GenBank Accession No. XM\_005591) and human TGF $\beta$  receptor type II mRNA (GenBank Accession No. XM 003094) are described in the art.

Cytokines of the TGF $\beta$  superfamily bind to specific serine/threonine kinase receptors and transmit intracellular signals through Smad proteins. Upon ligand stimulation, Smads move into the nucleus and function as components of transcription complexes. TGF $\beta$  signaling is regulated positively and negatively though various mechanisms. Positive regulation amplifies signals to a level sufficient for biological activity. Negative regulation occurs at the extracellular, membrane, cytoplasmic and nuclear levels.

Many tumors, including NSCLC and SCLC, produce high levels of active TGF $\beta$  (Constam et al. 1992; Eastham et al. 1995; Friedman et al. 1995; Jakowlew et al. 1995; Kong et al. 1995; Yamada et al. 1995; Eder et al. 1996). Elevated TGF $\beta$  levels have also been linked with immunosuppression (Sporn et al. 1986; Massague 1987; Bodmer et al. 1989; Border and Rouslahti 1992; Chen et al. 1997). TGF $\beta$  inhibits T cell activation in response to antigen stimulation. Additionally, TGF $\beta$  has antagonistic effects on the Natural Killer (NK) cells as well as the induction and proliferation of the lymphokine-activated killer (LAK) cells (Rook et al. 1986; Kasid et al. 1988; Tsunawaki et al. 1988; Hirte et al. 1991; Ruffini et al. 1993; Naganuma et al. 1996). In support of this, a relationship between TGF $\beta$  levels and survival has been demonstrated in colon cancer (Friedman et al. 1995). Recurrence rates were 18 fold higher in patients whose tumor produced high levels of TGF $\beta$  compared to those whose tumor produced low levels. This relationship was independent of nodal status and the degree of differentiation of the primary tumor.

Given the role of TGF $\beta$  in immune suppression we set out to evaluate the effect of TGF $\beta$  inhibition by NSCLC tumor vaccination. Using a TGF $\beta$  inhibitor approach we transfected a number of NSCLC cells with TGF $\beta$  antisense, selected from TGF $\beta$ 1, TGF $\beta$ 2, and TGF $\beta$ 3, and mixtures thereof. TGF $\beta$ 2 antisense was chosen as it demonstrated superiority in downregulating TGF $\beta$  expression compared with TGF $\beta$ 1 or a combination of TGF $\beta$ 1 and TGF $\beta$ 2. These

genetically modified NSCLC cells were then irradiated to prevent proliferation and were injected into a number of different animal tumor subjects. We observed that NSCLC cells previously ineffective as a component of a vaccine could be rendered efficacious through such a genetic modification. Blocking  $TGF\beta$  expression increased the immunogenicity of these animals. Furthermore such vaccinations eradicated previously implanted tumors and protected animals from tumor challenge.

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Given the role of  $TGF\beta$  in immune suppression we envision evaluating the effect of  $TGF\beta$  inhibition by SCLC tumor vaccination. Using a  $TGF\beta$  inhibitor approach we envision transfecting a number of SCLC cells with  $TGF\beta$  antisense, selected from  $TGF\beta1$ ,  $TGF\beta2$ , and  $TGF\beta3$ , and mixtures thereof. These genetically modified SCLC cells are then irradiated to prevent proliferation and are injected into a number of different animal tumor subjects. We envision observing that SCLC cells previously ineffective as a component of a vaccine would be rendered efficacious through such a genetic modification. Blocking  $TGF\beta$  expression is envisioned as increasing the immunogenicity of these animals. Furthermore such vaccinations are envisioned as eradicating previously implanted tumors and protecting animals from tumor challenge.

We have shown the efficacy of this approach in a NSCLC tumor model. In the KLN-205 NSCLC tumor model, DB2 mice were vaccinated with two injections of 5 x  $10^5$  irradiated TGF $\beta$ 2 antisense gene modified autologous NSCLC cells. This was capable of protecting the animals against a subsequent intraperitoneal (i.p.) tumor challenge with  $10^6$  unmodified KLN-205 NSCLC cells. In eradication experiments in this lung cancer tumor model, vaccination of animals bearing one week old tumors, with TGF $\beta$ 2 antisense gene modified cells resulted in marked tumor regression and prolonged tumor free survival compared to the control group.

Fakhrai et al. 1996 demonstrated the efficacy of this approach in a rat glial tumor. In the 9L gliosarcoma tumor model, intracranial implantation of as few as 300 tumor cells in Fisher-344 rat resulted in over 99% fatality after six weeks. Fakhrai et al. 1996 implanted 5 x  $10^3$  tumor cells into the brain of rats and administered tumor vaccinations. Animals immunized with TGF $\beta$ 2 antisense modified 9L cells, or with TGF $\beta$ 2 antisense modified 9L cells genetically modified to secrete IL-2 remained tumor free for the duration of the study (24 out of 24 or 100% tumor free survival). In contrast, the majority of the control group (2 out of 15) immunized with cells containing the empty vector developed tumors and had to be euthanized within five weeks (13% tumor free survival, p < 0.01).

Liau et al. 1998 demonstrated comparable efficacy of TGF $\beta$ 2 antisense gene therapy in a rat C-6 glioma tumor model. Dorigo et al. 1998 showed the efficacy of this approach in a murine ovarian teratoma (MOT) tumor model; however, only the group inoculated with TGF $\beta$  antisense and IL-2 gene modified cells resulted in significant protection from a subsequent tumor challenge, thus establishing the empiricism of the approach. Other groups have demonstrated similar anti-tumor effects of TGF $\beta$  gene therapy in cultured cells and animal tumor models (Kim et al. 1997).

Gene therapy has received considerable attention in recent years. Vaccination with tumor cells designed to augment tumor antigen presentation and induce specific anti-tumor immunity has yielded promising but limited results.

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Advances in our understanding of cancer biology and developments in vector technologies are advancing the therapeutic potential of tumor vaccination. It is now possible to genetically modify tumor cells for vaccination to express specific tumor suppressor genes, immune modulators, drug sensitive genes or antisense gene fragments (Huber et al. 1991; Culver et al. 1992; Trojan et al. 1992; Dranoff et al. 1993; Ram et al. 1993; Trojan et al. 1993).

A number of clinical studies have evaluated genetically modified allogeneic tumor cell as primary components of immunotherapeutic treatments for brain, skin, colon and breast cancers. The vaccination regimens have been shown to be safe and to generate humoral and cellular anti-vaccine immune responses. Preliminary results from several phase I clinical trials using gene therapies in patients with NSCLC have also demonstrated the safety of gene therapies approaches for this patient population (Dubinett, 1998; Roth, 1998; Swisher et al. 1998) as well as a SCLC patient population.

Groups have demonstrated experience in the field of gene therapy, both in a number of animal tumor models and in the clinic. (E.g., Fakhrai et al. 1995; Sobol et al. 1999.) The FDA has previously approved at least four INDs that have been submitted investigating gene-modified vaccination in patients with cancer:

- Sobol et al., BB-IND # 5812: "Injection of colon carcinoma patients with autologous irradiated tumor cells and fibroblasts genetically modified to secrete interleukin-2 (IL-2). A Phase I study"
- Sobol et al., BB-IND # 4840: "Active immunotherapy of glioblastoma with tumor cells or fibroblasts genetically modified to secrete interleukin-2 (IL-2)"
- Sobol et al., BB-IND # 7483: "A Phase I Study of Allogeneic Tumor Cells Genetically Modified to Express B7.1
   (CD80) Mixed with Allogeneic Fibroblasts Genetically Modified to Secrete IL-2 in Patients with Colorectal Carcinoma"
  - Fakhrai et al. BB-IND # 6658: "Proposal for a Phase I Clinical Trial: A Phase I Study of the Safety of Injecting
     Malignant Glioma Patients with Irradiated TGFβ2 Antisense Gene-Modified Autologous Tumor Cells"

In BB-IND #6658, the FDA approved a Phase I IND evaluating TGF $\beta$ 2 antisense gene therapy in patients with high grade glioma. Patients were vaccinated with autologous glioma tumor cells genetically modified with the a TGF $\beta$ 2 antisense plasmid to block TGF $\beta$ 2 expression. Therapy consisted of intradermal injections with 5 x 10<sup>6</sup>, 1 x 10<sup>7</sup> or 2 x 10<sup>7</sup> cells every 3 weeks for the first 4 months and every 1-2 months thereafter. To date, 5 patients have been treated. Under the same IND, the FDA approved the compassionate use of a partially haplotype matched allogeneic glioma cell line which was gene modified with the same TGF $\beta$ 2 antisense vector in a patient with pediatric glioma.

Overall treatment has been well tolerated with only low grade, transient toxicities reported. No significant adverse reactions at the immunization sites and no treatment related abnormalities have been observed on monitoring of complete blood counts, serum chemistries and urinalyses. In a few cases transient, mild erythema has been observed at the injection sites following the second and third subcutaneous injections with  $TGF\beta2$  antisense genemodified autologous tumor cells.

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Increased levels of CD3+, CD4+ and CD8+ effector cell infiltrates at the injection site and in secondary tumor biopsies have been observed. Immune histology of injection site biopsies and tumor obtained at subsequent operation demonstrates significantly higher number of immune infiltrates in comparison to biopsies taken prior to initiation of gene therapy.

Of the 5 patients treated, 1 patient demonstrated a clinical response, 2 demonstrated enhanced immune response, 1 showed tumor progression while the fifth patient is still undergoing therapy. In the patient who had a clinical response, overall MRI scans performed at approximately 6-week intervals during the first three months of treatment revealed modest changes in overall tumor size. Waxing and waning of peri-tumoral edema associated with alterations in Decadron doses could be observed. However, MRI scans showed tumor regression by seven months with a further improvement in response 3 months later.

The phase I clinical trial thus demonstrated the safety of injecting patients with  $5 \times 10^8$ ,  $1 \times 10^7$  or  $2 \times 10^7$  of TGF $\beta$ 2 antisense gene-modified autologous or haplotype-matched tumor cells. Furthermore, it is encouraging to see enhanced immunogenicity and preliminary clinical responses seen with this vaccination regimen. We contemplate the application of this gene therapy approach in patients with NSCLC or SCLC.

The invention encompasses methods and compositions for prolonging the survival of a subject having a non-small cell lung cancer (NSCLC) or a small cell lung cancer (SCLC) comprising administering to the subject a therapeutically effective amount of genetically modified cells containing a genetic construct expressing a  $TGF\beta$  inhibitor effective to reduce expression of  $TGF\beta$ , where the genetically modified cells are NSCLC or SCLCcells. Any method which neutralizes  $TGF\beta$  or inhibits expression of the  $TGF\beta$  gene (either transcription or translation) can be used to effectuate subject survival. Such approaches can also be useful for treatment applications, i.e., to treat NSCLC or SCLC.

In one embodiment, survival modalities can be designed to reduce the level of endogenous TGF $\beta$  gene expression, e.g., using antisense or ribozyme approaches to reduce or inhibit translation of TGF $\beta$  mRNA transcripts; triple helix approaches to inhibit transcription of the TGF $\beta$  gene; or targeted homologous recombination to inactivate or "knock out" the TGF $\beta$  gene or its endogenous promoter.

Antisense approaches involve the design of oligonucleotides (either DNA or RNA) that are complementary to TGF $\beta$  mRNA. The antisense oligonucleotides will bind to the complementary TGF $\beta$  mRNA transcripts and prevent translation. Absolute complementarity, although preferred, is not required. A sequence "complementary" to a portion of an RNA, as referred to herein, means a sequence having sufficient complementarity to be able to hybridize with the RNA, forming a stable duplex; in the case of double-stranded antisense nucleic acids, a single strand of the duplex DNA may thus be tested, or triplex formation may be assayed. The ability to hybridize will depend on both the degree of complementarity and the length of the antisense nucleic acid. Generally, the longer the hybridizing nucleic acid, the more base mismatches with an RNA it may contain and still form a stable duplex (or triplex, as the case may be). One

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skilled in the art can ascertain a tolerable degree of mismatch by use of standard procedures to determine the melting point of the hybridized complex.

Oligonucleotides that are complementary to the 5' end of the message, e.g., the 5' untranslated sequence up to and including the AUG initiation codon, should work most efficiently at inhibiting translation. However, sequences complementary to the 3' untranslated sequences of mRNAs have been shown to be effective at inhibiting translation of mRNAs as well. See generally, Wagner, R., 1994, Nature 372:333-335. Thus, oligonucleotides complementary to either the 5'- or 3'- non-translated, non-coding regions of TGF $\beta$  could be used in an antisense approach to inhibit translation of endogenous TGF $\beta$  mRNA. Oligonucleotides complementary to the 5' untranslated region of the mRNA should include the complement of the AUG start codon. Antisense oligonucleotides complementary to mRNA coding regions could also be used in accordance with the invention. Whether designed to hybridize to the 5'-, 3'- or coding region of TGF $\beta$  mRNA, antisense nucleic acids should be at least six nucleotides in length, and are preferably oligonucleotides ranging from 6 to about 50 nucleotides in length. In specific aspects the oligonucleotide is at least 17 nucleotides, at least 25 nucleotides or at least 50 nucleotides.

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Regardless of the choice of target sequence, it is preferred that in vitro studies are first performed to quantitate the ability of the antisense oligonucleotide to inhibit gene expression. It is preferred that these studies utilize controls that distinguish between antisense gene inhibition and nonspecific biological effects of oligonucleotides. It is also preferred that these studies compare levels of the target RNA or protein with that of an internal control RNA or protein. Additionally, it is envisioned that results obtained using the antisense oligonucleotide are compared with those obtained using a control oligonucleotide. It is preferred that the control oligonucleotide is of approximately the same length as the test oligonucleotide and that the nucleotide sequence of the oligonucleotide differs from the antisense sequence no more than is necessary to prevent specific hybridization to the target sequence.

The oligonucleotides can be DNA or RNA or chimeric mixtures or derivatives or modified versions thereof, single-stranded or double-stranded. The oligonucleotide can be modified at the base moiety, sugar moiety, or phosphate backbone, for example, to improve stability of the molecule, hybridization, etc. The oligonucleotide may include other appended groups such as peptides (e.g., for targeting host cell receptors), or agents facilitating transport across the cell membrane (see, e.g., Letsinger et al., 1989, Proc. Natl. Acad. Sci. U.S.A. 86:6553-6556; Lemaitre et al., 1987, Proc. Natl. Acad. Sci. 84:648-652; PCT Publication No. W088/09810, published Dec. 15, 1988).

The antisense oligonucleotide may comprise at least one modified base moiety which is selected from the group including but not limited to 5-fluorouracil, 5-bromouracil, 5-chlorouracil, 5-iodouracil, hypoxanthine, xantine, 4-acetylcytosine, 5-(carboxyhydroxylmethyl) uracil, 5-carboxymethylaminomethyl-2-thiouridine, 5-carboxymethylaminomethyluracil, dihydrouracil, beta-D-galactosylqueosine, inosine, N6-isopentenyladenine, 1-methylguanine, 1-methylguanine, 2,2-dimethylguanine, 2-methyladenine, 2-methylguanine, 3-methylcytosine, 5-methylcytosine, N6-adenine, 7-methylguanine, 5-methylaminomethyluracil, 5-methoxyaminomethyl-2-thiouracil, beta-D-mannosylqueosine, 5'-methoxycarboxymethyluracil, 5-methoxyuracil, 2-methylthio-N6-isopentenyladenine, uracil-5-oxyacetic acid (v), wybutoxosine, pseudouracil, queosine, 2-thiocytosine, 5-methyl-2-thiouracil, 2-thiouracil, 4-

thiouracil, 5-methyluracil, uracil5-oxyacetic acid methylester, uracil-5-oxyacetic acid (v), 5-methyl-2-thiouracil, 3-(3-amino-3-N-2-carboxypropyl) uracil, (acp3)w, and 2,6-diaminopurine.

The antisense oligonucleotide may also comprise at least one modified sugar moiety selected from the group including but not limited to arabinose, 2-fluoroarabinose, xylulose, and hexose.

In yet another embodiment, the antisense oligonucleotide comprises at least one modified phosphate backbone selected from the group consisting of a phosphorothioate, a phosphorodithioate, a phosphoramidate, a phosphordiamidate, a methylphosphonate, an alkyl phosphotriester, and a formacetal or analog thereof.

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Oligonucleotides of the invention may be synthesized by standard methods known in the art, e.g. by use of an automated DNA synthesizer (such as are commercially available from Biosearch, Applied Biosystems, etc.). As examples, phosphorothicate oligonucleotides may be synthesized by the method of Stein et al. (1988, Nucl. Acids Res. 16:3209), methylphosphonate oligonucleotides can be prepared by use of controlled pore glass polymer supports (Sarin et al., 1988, Proc. Natl. Acad. Sci. U.S.A. 85:7448-7451), etc.

However, it is often difficult to achieve intracellular concentrations of the antisense sufficient to suppress translation of endogenous mRNAs. Therefore a convenient approach utilizes a recombinant DNA construct in which the antisense sequence is placed under the control of a strong promoter. The use of such a construct to transfect target cells will result in the transcription of sufficient amounts of single stranded RNAs that will form complementary base pairs with the endogenous TGFβ transcripts and thereby prevent translation of the TGFβ mRNA. For example, a vector can be introduced such that it is taken up by a cell and directs the transcription of an antisense RNA. Such a vector can remain episomal or become chromosomally integrated, as long as it can be transcribed to produce the desired antisense RNA. Such vectors can be constructed by recombinant DNA technology methods standard in the art. Vectors can be plasmid, viral, or others known in the art, used for replication and expression in mammalian cells. Expression of the sequence encoding the antisense RNA can be by any promoter known in the art to act in mammalian, preferably human cells. Such promoters can be inducible or constitutive. Such promoters include but are not limited to: the SV40 early promoter region (Bernoist and Chambon, 1981, Nature 290:304-310), the promoter contained in the 3' long terminal repeat of Rous sarcoma virus (Yamamoto et al., 1980, Cell 22:787-797), the herpes thymidine kinase promoter (Wagner et al., 1981, Proc. Natl. Acad. Sci. U.S.A. 78:1441-1445), the regulatory sequences of the metallothionein gene (Brinster et al., 1982, Nature 296:39-42), etc.

Ribozyme molecules-designed to catalytically cleave TGF $\beta$  mRNA transcripts can also be used to prevent translation of TGF $\beta$  mRNA and expression of TGF $\beta$ . (See, e.g., PCT International Publication W090/11364, published Oct. 4, 1990; Sarver et al., 1990, Science 247:1222-1225). While ribozymes that cleave mRNA at site specific recognition sequences can be used to destroy TGF $\beta$  mRNAs, the use of hammerhead ribozymes is preferred. Hammerhead ribozymes cleave mRNAs at locations dictated by flanking regions that form complementary base pairs with the target mRNA. The sole requirement is that the target mRNA have the following sequence of two bases: 5'-UG-3'. The construction and production of hammerhead ribozymes is well known in the art and is described more fully

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in Haseloff and Gerlach, 1988, Nature, 334:585-591. There are hundreds of potential hammerhead ribozyme cleavage sites within the nucleotide sequence of a TGF $\beta$  cDNA. Preferably the ribozyme is engineered so that the cleavage recognition site is located near the 5' tend of the TGF $\beta$  mRNA; i.e., to increase efficiency and minimize the intracellular accumulation of non-functional mRNA transcripts.

The ribozymes of the present invention also include RNA endoribonucleases (hereinafter "Cech-type ribozymes") such as the one which occurs naturally in Tetrahymena Thermophila (known as the IVS, or L-19 IVS RNA) and which has been extensively described by Thomas Cech and collaborators (Zaug, et al., 1984, Science, 224:574-578; Zaug and Cech, 1986, Science, 231:470-475; Zaug, et al., 1986, Nature, 324:429-433; published International patent-application No. WO 88/04300 by University Patents Inc.; Been and Cech, 1986, Cell, 47:207-216). The Cechtype ribozymes have an eight base pair active site which hybridizes to a target RNA sequence whereafter cleavage of the target RNA takes place. The invention encompasses those Cech-type ribozymes which target eight base-pair active site sequences that are present in TGFβ.

As in the antisense approach, the ribozymes can be composed of modified oligonucleotides (e.g. for improved stability, targeting, etc.) and need be delivered to target cells which express TGF $\beta$ . A convenient method of delivery involves using a DNA construct encoding the ribozyme under the control of a strong promoter so that transfected cells will produce sufficient quantities of the ribozyme to destroy endogenous TGF $\beta$  messages and inhibit translation. Because ribozymes unlike antisense molecules, are catalytic, a lower intracellular concentration is required for efficiency.

Endogenous TGF $\beta$  gene expression can also be reduced by inactivating or "knocking out" the TGF $\beta$  gene or its promoter using targeted homologous recombination. (E.g., see Smithies et al., 1985, Nature 317:230-234; Thomas & Capecchi, 1987, Cell 51:503-512; Thompson et al., 1989 Cell 5:313-321). For example, a mutant, non-functional TGF $\beta$  (or a completely unrelated DNA sequence) flanked by DNA homologous to the endogenous TGF $\beta$  gene (either the coding regions or regulatory regions of the TGF $\beta$  gene) can be used, with or without a selectable marker and/or a negative selectable marker, to transfect target cells that express TGF $\beta$ . Insertion of the DNA construct, via targeted homologous recombination, results in inactivation of the TGF $\beta$  gene.

Alternatively, endogenous TGF $\beta$  gene expression can be reduced by targeting deoxyribonucleotide sequences complementary to the regulatory region of the TGF $\beta$  gene (i.e., the TGF $\beta$  promoter and/or enhancers) to form triple helical structures that prevent transcription of the TGF $\beta$  gene in target cells. (See generally, Helene, C. 1991, Anticancer Drug Des., 6(6):569-84; Helene, C., et al., 1992, Ann, N.Y. Accad. Sci., 660:27-36; and Maher, L. J., 1992, Bioassays 14(12):807-15).

In yet another embodiment of the invention, the activity of TGF $\beta$  can be reduced using a "dominant negative" approach to effectuate subject survival. To this end, genetic constructs which encode defective TGF $\beta$ s can be used to diminish the activity of TGF $\beta$  on neighboring cells. For example, nucleotide sequences that direct expression of TGF $\beta$ s in which domains are deleted or mutated can be introduced into target cells. Alternatively, targeted homologous

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recombination can be utilized to introduce such deletions or mutations into the target cell's endogenous  $TGF\beta$  gene. The engineered cells will express non-functional cytokines (i.e., a cytokine that is capable of binding its natural receptor, but incapable of signal transduction). Such engineered cells should facilitate a diminished response on neighboring cells to endogenous  $TGF\beta$  ligand, resulting in subject survival.

In an alternative embodiment, the administration of genetic constructs encoding soluble peptides, proteins, fusion proteins, or antibodies that bind to and "neutralize" intracellular TGF $\beta$  effectuate subject survival. To this end, genetic constructs encoding peptides corresponding to domains of the TGF $\beta$  receptor, deletion mutants of the TGF $\beta$  receptor, or either of these TGF $\beta$  receptor domains or mutants fused to another polypeptide (e.g., an IgFc polypeptide) can be utilized. Alternatively, genetic constructs encoding anti-idiotypic antibodies or Fab fragments of antiidiotypic antibodies that mimic the TGF $\beta$  receptor and neutralize TGF $\beta$  can be used. Such genetic constructs encoding these TGF $\beta$  receptor peptides, proteins, fusion proteins, anti-idiotypic antibodies or Fabs are administered to neutralize TGF $\beta$  and effectuate subject survival.

Genetic constructs encoding antibodies that specifically recognize one or more epitopes of  $TGF\beta$ , or epitopes of conserved variants of  $TGF\beta$ , or peptide fragments of  $TGF\beta$  are also encompassed by the invention. Such antibodies include but are not limited to polyclonal antibodies, monoclonal antibodies (mAbs), humanized or chimeric antibodies, single chain antibodies, Fab fragments, F(ab')2 fragments, fragments produced by a Fab expression library, and epitope-binding fragments of any of the above. Genetic constructs encoding such antibodies may be used as a method for the inhibition of  $TGF\beta$  activity and effectuation of subject survival.

For the production of antibodies, various host animals may be immunized by injection with TGF $\beta$ , a TGF $\beta$  peptide, truncated TGF $\beta$ , functional equivalents of TGF $\beta$  or mutants of TGF $\beta$ . Such host animals may include but are not limited to rabbits, mice, and rats, to name but a few. Polyclonal antibodies are heterogeneous populations of antibody molecules derived from the sera of the immunized animals.

Monoclonal antibodies, which are homogeneous populations of antibodies to a particular antigen, may be obtained by any technique which provides for the production of antibody molecules by continuous cell lines in culture. These include, but are not limited to, the hybridoma technique of Kohler and Milstein, (1975, Nature 256:495-497; and U.S. Pat. No. 4,376,110), the human B-cell hybridoma technique (Kosbor et al., 1983, Immunology Today 4:72; Cole et al., 1983, Proc. Natl. Acad. Sci. USA 80:2026-2030), and the EBV-hybridoma technique (Cole et al., 1985, Monoclonal Antibodies And Cancer Therapy, Alan R. Liss, Inc., pp. 77-96). Such antibodies may be of any immunoglobulin class including IgG, IgM, IgE, IgA, IgD and any subclass thereof.

In addition, techniques developed for the production of "chimeric antibodies" (Morrison et al., 1984, Proc. Natl. Acad. Sci., 81:6851-6855; Neuberger et al., 1984, Nature, 312:604-608; Takeda et al., 1985, Nature, 314:452-454) by splicing the genes from a mouse antibody molecule of appropriate antigen specificity together with genes from a human antibody molecule of appropriate biological activity can be used. A chimeric antibody is a molecule in which

different portions are derived from different animal species, such as those having a variable region derived from a murine mAb and a human immunoglobulin constant region.

Alternatively, techniques described for the production of single chain antibodies (U.S. Pat. No. 4,946,778; Bird, 1988, Science 242:423-426; Huston et al., 1988, Proc. Natl. Acad. Sci. USA 85:5879-5883; and Ward et al., 1989, Nature 334:544-546) can be adapted to produce single chain antibodies against TGFβ gene products. Single chain antibodies are formed by linking the heavy and light chain fragments of the Fv region via an amino acid bridge, resulting in a single chain polypeptide.

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Antibody fragments which recognize specific epitopes may be generated by known techniques. For example, Fab expression libraries may be constructed (Huse et al., 1989, Science, 246:1275-1281) to allow rapid and easy identification of monoclonal Fab fragments with the desired specificity.

Additionally, the enzymes which cleave TGF $\beta$  precursors to the active isoforms may be inhibited in order to block activation of TGF $\beta$ . TGF $\beta$  must be activated to exhibit its biological effects and enzymes are required to cleave the precursor protein. These enzymes may be altered genetically to prevent interaction with the precursor protein, preventing cleavage of the protein to its mature form. Transcription or translation of these enzymes may be blocked by a means known to the art. These enzymes may alternatively be inhibited by any means known to one of skill in the art.

TGF $\beta$ s bind to serine/threonine kinase receptors and transmit intracellular signals through Smad proteins. Signal transduction may be interrupted in order to repress signaling initiated by TGF $\beta$ . By disrupting signal transduction it is possible to prevent the immunosuppressive effect of TGF $\beta$ . This may be accomplished by any means known in the art in which the interaction between the TGF $\beta$  receptor and the Smad protein is antagonized or prevented, including administering genetically modified cells which express proteins that block or compete with TGF $\beta$  receptor and Smad protein interactions. Alternatively, the transcription or translation of TGF $\beta$  receptor or Smad protein may be altered by any means known in the art in order to prevent signal transmission along the signaling pathway.

Target cells genetically engineered to express such a form of  $TGF\beta$  inhibitor are administered as an immunogen to patients with NSCLC or SCLC, whereupon they will serve to enhance anti-tumor immune responses and thereby prolong survival of tumor-bearing subjects. Such cells may be obtained from the patient (autologous) or a donor (allogeneic or xenogeneic). For a patient with NSCLC, the genetically engineered cells constitute non-small cell lung cancer (NSCLC) cells, which are NSCLC cells by virtue of being derived from a NSCLC or mimicing a NSCLC (i.e., having shared common tumor antigens or epitopes with a primary NSCLC). Alternatively, for a patient with SCLC, the genetically engineered cells constitute small cell lung cancer (SCLC) cells, which are SCLC cells by virtue of being derived from a SCLC or mimicing a SCLC (i.e., having shared common tumor antigens or epitopes with a primary SCLC).

Autologous cells are cell that are derived from the same individual. Allogeneic cells are cells that are derived from another individual of the same species so that the cells have intraspecies genetic variations. Xenogeneic cells are cells that are derived from an individual of a different species so that the cells have interspecies antigenic differences.

In one embodiment, an allogeneic (or xenogeneic) NSCLC or SCLC tumor cell line is chosen as the immunogen. Lung tumor cell lines have been shown to have shared epitopes with primary tumors (Takenoyama et al. 1998). These investigators showed that MHC class I restricted CTL generated against a human lung adenocarcinoma cell line had demonstrable cytotoxicity against another lung tumor cell line. The cross reactivity in these experiments was blocked by anti-MHC class I and anti-CD8 monoclonal antibodies, suggesting that shared common tumor antigens exist among lung cancer cells.

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In another embodiment, an allogeneic (or xenogeneic) cell cocktail is used as an immunogen in patients with NSCLC or SCLC. One can employ more than one, e.g., two, three, four or more cell lines rather than one to increase the total number of tumor antigens present.

In addition, the target cells will have low levels of  $TGF\beta$  expression owing to transfection with a genetic construct encoding a  $TGF\beta2$  inhibitor. Suppression of  $TGF\beta$  expression by the tumor cells will remove a major source of immune suppression operative at the site of vaccine injection. A local immune response, directed against the injected tumor cells will induce a systemic immune response against the patients' native tumor.

The target cells are genetically engineered in vitro using recombinant DNA techniques to introduce the genetic constructs into the cells, e.g., by transduction (using viral vectors) or transfection procedures, including but not limited to the use of plasmids, cosmids, YACs, electroporation, liposomes, etc. The engineered cells can be introduced into the patient, e.g., in the circulation, intraperitoneally, intradermally, subcutaneously, at the lobes of the lung. Alternatively, the cells can be incorporated into a matrix and implanted in the body as part of a tissue graft.

In another embodiment, target cells are engineered to express a coding sequence for one or more cytokines. In one alternative, an expression vector singly encoding the one or more cytokines is introduced into the target cells. In another alternative, an expression vector doubly encoding the one or more cytokines and a  $TGF\beta$  inhibitor is introduced into the target cells. In still another alternative, some target cells are engineered to express a coding sequence for one or more cytokines and other target cells are genetically modified to express a  $TGF\beta$  inhibitor. By co-administering the immunostimulatory agent along with inhibiting the immunosuppressant  $TGF\beta$ , a subject's immune response to tumor cells may be improved. Examples of cytokines useful for practice of the present invention include interleukin-1, interleukin-2, interleukin-3, interleukin-5, interleukin-6, interleukin-7, interleukin-8, interleukin-9, interleukin-10, interleukin-11, interleukin-12, interleukin-15, interferon-alpha, interferon-gamma, tumor necrosis factor-alpha, transforming growth factor-beta, granulocyte macrophage colony stimulating factor, and granulocyte colony stimulating factor. The level of cytokine expression should be regulated such that anti-tumor immunity can be increased without producing significant systemic toxicity in the subject.

When the target cells to be administered are non-autologous cells, they can be administered using well known techniques which prevent the development of a host immune response against the introduced cells. For example, the

cells may be introduced in an encapsulated form which, while allowing for an exchange of components with the immediate extracellular environment, does not allow the introduced cells to be recognized by the host immune system.

Toxicity and therapeutic efficacy of NSCLC or SCLC cells can be determined by standard pharmaceutical procedures in cell cultures or experimental animals, e.g., for determining the LD50 (the dose lethal to 50% of the population) and the ED50 (the dose therapeutically effective in 50% of the population). The dose ratio between toxic and therapeutic effects is the therapeutic index and it can be expressed as the ratio LD50 /ED50. Numbers of NSCLC or SCLC cells which exhibit large therapeutic indices are preferred.

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The data obtained from the cell culture assays and animal studies can be used in formulating a range of dosage for use in humans. The dosage for humans lies preferably within a range of concentrations that include the ED50 with little or no toxicity. For any number of NSCLC or SCLC cells used in the invention, the therapeutically effective dose can be estimated initially from cell culture assays. A dose may be formulated in animal models to achieve a concentration range that includes the IC50 (i.e., the concentration of the test material which achieves a half-maximal inhibition of symptoms) as determined in cell culture. Such information can be used to more accurately determine useful doses in humans.

Pharmaceutical compositions for use in accordance with the present invention may be formulated in a conventional manner using one or more physiologically acceptable carriers or excipients. Various adjuvants may be used to increase the immunological response, including QS-21, Detox-PC, MPL-SE, MoGM-CSF, TiterMax-G, CRL-1005, GERBU, TERamide, PSC97B, Adjumer, PG-026, GSK-1, GcMAF, B-alethine, MPC-026, Adjuvax, CpG ODN, Betafectin, Alum, and MF59 (see Kim et al. Vaccine 2000, 18: 597 and references therein). Formulations for injections may be presented in unit dosage form, e.g., in ampoules or multi-dose containers.

The present invention further provides a therapeutic composition comprising the genetically modified cells expressing a  $TGF\beta$  inhibitor and a therapeutically acceptable carrier. As used herein, a therapeutically acceptable carrier includes any and all solvents, including water, dispersion media, culture from cell media, isotonic agents and the like that are non-toxic to the host. Conveniently, it is an aqueous isotonic buffered solution with a pH of around 7.0. The use of such media and agents in therapeutic compositions is well known in the art. Except insofar as any conventional media or agent is incompatible with the genetically modified cells of the present invention, use of such conventional media or agent in the therapeutic compositions is contemplated. Supplementary active ingredients can also be incorporated into the compositions.

The therapeutic compositions of the present invention may be administered to an animal in need thereof. Accordingly, the present invention provides methods for inducing an immune response in an animal in need of such response, which comprise administering to an animal an immunologically effective amount of the subject genetically modified cells. The present invention also provides methods for preventing or treating a tumor in an animal, which comprise administering to an animal an anti-tumor effective amount of the subject genetically modified cells.

The term "animal" used herein encompasses all mammals, including human. Preferably, the animal of the present invention is a human subject.

The immune response induced in the animal by administering the subject genetically modified cells may include cellular immune responses mediated primarily by cytotoxic T cells, capable of killing tumor cells, as well as humoral immune responses mediated primarily by helper T cells, capable of activating B cells thus leading to antibody production. A variety of techniques may be used for analyzing the type of immune responses induced by the subject genetically modified cells, which are well described in the art; e.g., Coligan et al. Current Protocols in Immunology, John Wiley & Sons Inc. (1994).

The term "preventing a tumor" used herein means the occurrence of the tumor is prevented or the onset of the tumor is significantly delayed. The term "treating a tumor" used herein means that the tumor growth is significantly inhibited, which is reflected by, e.g., the tumor volume. Tumor volume may be determined by various known procedures, e.g., obtaining two dimensional measurements with a dial caliper.

When "an immunologically effective amount", "an anti-tumor effective amount", or "a tumor-inhibiting effective amount" is indicated, the precise amount of the genetically modified cells to be administered can be determined by a physician with consideration of individual differences in age, weight, tumor size, extent of infection or metastasis, and condition of the patient. It can generally be stated that a therapeutic composition comprising the subject genetically modified cells is conveniently administered in an amount of at least about 1 times 10<sup>3</sup> to about 5 times 10<sup>9</sup> cells per dose.

The administration of the subject therapeutic compositions may be carried out in any convenient manner, including by aerosol inhalation, injection, ingestion, transfusion, implantation or transplantation. Conveniently, the genetically modified cells of the present invention are administered to a patient by subcutaneous (s.c.), intraperitoneal (i.p.), intra-arterial (i.a.), or intravenous (i.v.) injection. The therapeutically acceptable carrier should be sterilized by techniques known to those skilled in the art.

The invention is further illustrated by the following specific example which is not intended in any way to limit the scope of the invention. We envision substituting TGF $\beta$ 1 and TGF $\beta$ 3 for TGF $\beta$ 2 in the following example. Additionally, we envision substituting SCLC cells for NSCLC cells in the following example.

25 Example

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In a clinical trial we use four human non-small cell lung cancer cell lines that have been previously established in tissue culture laboratory. We gene modify these tumor cells in the laboratory to block their  $TGF\beta$  secretion. We then use the genetically engineered cells as vaccines in patients with non-small cell lung cancer. Patients are injected four times, in monthly intervals, with the gene modified vaccine cocktails that constitute the four non-self (allogeneic)  $TGF\beta$  antisense gene modified tumor cells. Our rationale for using other people's tumor cells is that lung tumor cell lines belonging to different people have been shown to share common characteristics that are recognized by non-self immune systems. Treated patients are evaluated four months after they enter therapy. Patients that respond to therapy receive an additional four to twelve injections to evaluate whether their response to therapy can be amplified.

Patients are randomly assigned to one of three separate cohorts. The vaccine cocktail constitutes an equal number of each of the four irradiated TGF $\beta$  antisense gene modified NSCLC cell lines. The number of injected cells in the three cohorts is  $1.25 \times 10^7$ ,  $2.5 \times 10^7$ , and  $5 \times 10^7$  cells respectively.

Response, time to tumor progression, and tumor free survival are monitored in patients and compared with historical controls and patients receiving other forms of therapy. Patients are monitored and evaluated according to standard evaluation criteria of no response, stable disease, partial response and complete response. The results of this study are used to evaluate the feasibility of additional clinical trial with  $TGF\beta$  antisense gene modified tumor cells alone and in combination with IL-2 (or other cytokine) gene modification.

# Primary Objective

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The primary objective of the clinical trial is to evaluate the ability of increasing doses of a gene-modified tumor cell vaccine to induce tumor response in patients with NSCLC.

# Study Design

This study is designed to evaluate the efficacy of immunization with increasing doses of an allogeneic tumor cell vaccine in patients with NSCLC. Patients are followed for clinical response, immunogenicity and safety.

Eligible patients receive 4 monthly intradermal injections with a cell cocktail comprised of equal numbers of four irradiated allogeneic TGF $\beta$ 2 antisense gene modified NSCLC cell lines. Patients are randomized to one of the three study cohorts. Patients receive 1.25 x 10<sup>7</sup>, 2.5 x 10<sup>7</sup>, or 5 x 10<sup>7</sup> gene modified cells respectively.

When available, tumor samples obtained from study patients at the time of clinically indicated surgery are used to establish a cell line for each patient. The patients' tumor cells are then used in precursor analyses or cytotoxicity assay monitoring of the patients immune responses to gene therapy inoculations.

#### Vaccine Administration

Patients receive intradermal injections of the tumor cell vaccine at months 0, 1, 2, 3 and 4. These are administered in an outpatient setting. The sites of injection are rotated between the upper and lower extremities. Patients are observed in the clinic for 2 hours following vaccination. During this observation period, vital signs are taken every 30 minutes. Patients experiencing no significant side effects from treatment are discharged 2 hours after vaccination.

#### **Outline of Study Procedures**

Patients are vaccinated according to the schedule outlined in the table below. Patients are initially treated once a month for 4 months, unless there is documented unmanageable toxicity or clinically significant disease progression requiring intervention with other anti-cancer therapies. Tumor staging (by comprehensive CT/MRI scans) is performed at baseline and at weeks 8, 16 and 28 and every 3 months thereafter. Patients have serial monitoring of immune response (humoral and T cell responses) every 4 weeks up to week 28 and every 12 weeks thereafter. Patients are monitored closely for toxicity throughout the study. In those patients demonstrating benefit from treatment, additional vaccinations, given every 4-8 weeks, are given for up to 12 additional vaccinations (total of 16 vaccinations).

# **Stopping Rules**

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Given that tumor responses from vaccination may follow a period of initial tumor progression, patients are allowed to stay on study in the face of non-clinically significant progression at week 8. At week 16, such patients must shown no further tumor progression (no more than a 25% increase at week 16 as compared to week 8). Patients who demonstrate progressive disease at week 16 (compared to week 8) are removed from study.

**Overview of Monthly Treatment Schema** 

Procedure	Screen	Day	Day	Day	Day
		1	2	8	29/1
Informed consent	Х	· · · · · ·			
History, Exam <sup>1</sup>	X [2]	Х			Х
Phone contact			Х		
Vital signs, weight, PS	X [2]	Х		Х	Х
Adverse events	X [2]	Х		Х	Х
Concomitant medications	X [2]	Х		Х	Х
CBC with differential	X [1]	Χ		Х	Х
Electrolyte panel	X [1]	Х			Х
Metabolism panel	X [2]	Х			Х
Tumor staging <sup>2</sup>	X [4]				
Biopsy of inoculation site				Х	
Humoral Immunity <sup>3</sup>	X [2]				Х
Cellular Immunity <sup>3</sup>	X [2]				Х
Vaccine administration <sup>4</sup>		Х			Х

[X]: within X weeks prior to day 1

10 <sup>3</sup> repeated at weeks 4, 8, 12, 16, 20, 24, 28, then every 12 weeks

Definition of procedures

Phone contact:

Patients are contacted by phone by the study nurse to assess the degree of inflammation, pain, or puritis at local injection site

once a month during treatment and once every 3 months in follow-up

<sup>&</sup>lt;sup>2</sup>repeated at weeks 8, 16, 28, then every 12 weeks

<sup>&</sup>lt;sup>4</sup>administered at weeks 1, 4, 8, 12. Continued administration possible for responding patients

CBC with differential: WBC, HCT, HGB, platelet count, % neutrophil, % lymphocytes, %

monocytes

Electrolyte panel: Sodium, potassium, chloride, carbon dioxide, BUN, creatinine, glucose

Metabolism panel: Calcium, phosphorus, AST, ALT, alkaline phosphatase, bilirubin, uric

acid, albumin, protein,

Tumor staging: Physical exam, x-rays, CT/MRI as appropriate. All staging should use

the same method to assess tumor as used at baseline

Biopsy of inoculation site: Punch skin biopsy performed at periphery of inflamed vaccination site

or, if no inflammation, to include vaccination site

10 Humoral Immunity: Serum anti-tumor titers

Cellular Immunity: Immunophenotyping of peripheral blood B-cell and T-cell subsets (if

sufficient cells are available) including CD3, CD4, CD8, CD16, CD20,

and CD68.

Measure PBC cytokine profile by quantitative (semi-quantitative), NK

activity (nonspecific killing), LAK activity (Allo-killing).

#### **Inclusion Criteria**

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Signed informed consent

≥ 18 years

- Histologically confirmed non-curable NSCLC with measurable disease and an estimated volume of 125 cc
- Performance status (ECOG)  $\leq 2$ 
  - Absolute granulocyte count 1,500/mm3
  - Platelet count 100,000/mm3
  - Total Bilirubin ≤ 2 mg/dL
  - AST and ALT ≤ 2x Upper Limit of Normal
- Creatinine ≤ 1.5 mg/dL

# **Exclusion Criteria**

Concurrent systemic steroids > 20 mg prednisone/day

- Prior splenectomy
- Surgery, chemotherapy, radiotherapy, steroid therapy or immunotherapy < 4 weeks of study entry</li>
- Brain metastases or meningeal lymphomatosis unless treated and stable for 2 months
- Known HIV positive
- Serious non-malignant disease (e.g., congestive heart failure, or active uncontrolled bacterial, viral, or fungal
  infections), or other conditions which, in the opinion of the investigator would compromise protocol objectives.
  - Prior malignancy (excluding nonmelanoma carcinomas of the skin) unless in remission for  $\geq 2$  years
  - Treatment with an investigational drug within 30 days prior to study entry
  - History of psychiatric disorder that would impede adherence to protocol
- Pregnant or nursing women or refusal to practice contraception if of reproductive potential

#### Conduct of the Study

The study is conducted according to Good Clinical Practice, the Declaration of Helsinki and US 21 CFR Part 50 - Protection of Human Subjects, and Part 56 - Institutional Review Boards. Written, dated informed consent for the study is obtained from all patients before protocol-specified procedures are carried out. After signing, patients are given a copy of their informed consent. Approval of this study is obtained from the appropriate Institutional Review Board prior to enrolling patients on study. Consent forms are in a language fully comprehensible to the prospective patient. Consent is documented either by the patient's dated signature or by the signature of an independent witness who records the patient's consent.

#### **Tumor Response**

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Patients are evaluated by CT/MRI and physical examination. Response is reported using standard outcome measures for clinical trials (complete response (CR), partial response (PR), stable disease (SD) and progressive disease (PD)). Any response to treatment (either PR or CR) requires two confirmatory staging at least 4 weeks apart.

#### Complete response

- Resolution of all measurable disease for a period of at least 4 weeks
- Resolution of all evaluable disease for a period of at least 4 weeks
  - No new lesions (either measurable or evaluable)

#### Partial response

Decrease in the sum of the product of all measurable lesions by at least 50% for a period of at least 4 weeks

Subjective improvement in evaluable disease for a period of at least 4 weeks

No new lesions (either measurable or evaluable)

Stable disease

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Less than a 50% decrease AND less than a 25% increase in the sum of the products of all measurable lesions

No new lesions (either measurable or evaluable)

Progressive disease

 Greater than a 25% increase in the sum of the products of all measurable lesions <u>OR</u> new lesions (either measurable or evaluable)

### 10 Evaluation of Immune Response

Humoral immune response assessment

Humoral anti-tumor immune responses is evaluated by comparing the titer of pre-treatment and post-treatment sera for reactivity against the vaccinating cell lines using an enzyme-linked immunosorbent assay (ELISA). Briefly, 10<sup>5</sup> target cells are immobilized on filter paper disks in a 96-well incubator chamber (V and P Enterprises, La Jolla, CA) and then incubated for 30 minutes with the test sera. The plates are washed and then incubated with an enzyme-conjugated anti-human lg. The plates are again washed, the enzyme substrate is added, and the binding is quantitated by measuring the absorbance of each well on an ELISA reader.

Cellular immune response assessment

Immunophenotyping

Standard immunofluorescence flow cytometry assays are performed to assess patients pre and post treatment immune effector cells profiles. Percentages of effector cell subpopulations reacting with monoclonal antibodies to T-cells (CD3, CD4, CD8), natural killer cells (CD16) and B-cells (CD20) are measured in the pre- and post-treatment peripheral blood lymphocyte population and correlated with patients responses measured by other criteria. Briefly, the Ficoll-Hypaque purified mononuclear cells are incubated with the primary antibody for 1 hour at room temperature, washed and then incubated with fluorochrome conjugated secondary antibody. The cells are washed, fixed, and the percentage of positive cells are determined with a flow cytometer. Incubations of the cells with isotype-matched control antibody instead of the primary antibody serve as negative controls.

Natural Killer (NK) activity

NK activity is analyzed using a standard chromium release assay using the NK-sensitive cell line K562 as the target. Briefly, K562 cells are labeled by incubating them with  $^{51}$ Cr for 45 minutes at 37°C. The target cells are washed extensively and then  $5 \times 10^3$  K562 are incubated for 4 hours at 37°C with pre- and post-treatment PBMC at effector cell:target cell ratios ranging from 100:1 to 3:1. The cells arel then centrifuged and the amount  $^{51}$ Cr-release is

measured using a gamma counter. The percent specific lysis is determined using the formula: (experimental cpm - background cpm)/(total cpm - background cpm)  $\times$  100.

LAK activity

LAK activity is determined by chromium release assay as described above, using the LAK-sensitive cell line DAUDI as the target.

Pre- and post-treatment cytokine profile of lymphocytes

The cytokine profile of the patients PBMC is determined by semi-quantitative PCR assays. RNA is extracted from patients pre- and post-treatment purified mononuclear cells and used to synthesize first strand cDNA by an Invitrogen (San Diego, CA) cDNA cycle kit according to the manufacturer's recommendation. The first strand cDNA is then used as template in PCR assays employing different primer sets for detection of IL-2, IL-4, IL-6, IL-7, IL-10, GM-CSF, γ-INF, TNF-α, etc. To achieve quantitation PCR reactions are limited to 15-18 cycles. As an internal control and to aid in quantitation of the products known concentrations of a control RNA are added to each sample prior to initiation of cDNA synthesis. Specific primers for the control sequence are then added to the PCR reactions. Patient samples cytokine profiles are determined by quantitating patients' PCR products and comparing them with the control PCR products.

Skin biopsy of immunization site

Standard hematoxylin and eosin staining and immunohistochemical methods employing monoclonal antibodies to hematopoietic cell subsets are employed to characterize the immune infiltrates observed in skin biopsies at immunization sites. Monoclonal antibodies to T-cells (CD3, CD4, CD8), natural killer cells (CD16) and B-cells (CD20) are utilized for these studies. Briefly, for the immunohistochemical studies, cryostat sections are fixed in cold acetone and then incubated with primary antibody for 1 hour at room temperature. The sections are washed and then incubated with horseradish peroxidase conjugated secondary antibody followed by staining sections with an appropriate chromagen substrate and examined by light microscopy. Incubations of sections with isotype-matched control antibody instead of the primary antibody serve as negative controls.

# 25 Drug Information

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**Clinical Formulation** 

The vaccine is provided in frozen vials containing at least 20 x 10<sup>8</sup> cells per vial.

**Pharmacists Instructions** 

**Undiluted** material

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Vial volume:

1 ml

Appearance:

Cloudy fluid

Storage:

-176 °C (Liquid Nitrogen).

Hazards:

Frozen vials are not considered hazardous if

unbroken. Vials contain cell frozen in a mixture containing 10% dimethyl sulphoxide and 50% fetal

calf serum.

Handling:

Frozen vials are not considered a safety hazard if

unbroken. Broken vials should be disposed of in

accordance with biohazard procedures for

cytotoxic drugs.

Diluted material

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10 Preparation: Before being injected into patients, a frozen vial is

thawed in a biosafety hood and washed twice

with serum containing medium and four times with

lactated Ringer's. The cells are then counted and

adjusted to the appropriate number of cells per

injection in a volume of 250-400  $\mu$ l. The cell

suspension is delivered in a capped 1mL syringe.

Drug concentrate:  $1.25 \times 10^7$ ,  $2.5 \times 10^7$ , or  $5 \times 10^7$  cells per injection

in a volume of 250-400  $\mu$ l.

Diluent: Lactated Ringer's

20 Route of administration: Intradermal injection

Storage

Frozen, unopened vials are stored at -176°C (Liquid Nitrogen).

# 25 Data Evaluation

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Statistics and Estimated Sample Size

Patients in the amount of 27-75 are enrolled. This is a two-stage study. Each of the three treatment arms initially accrues 9 patients. Should no responses be seen in the first 9 patients, then no further patients are accrued to that treatment arm. If at least 1 response is seen in the first 9 patients, then 16 additional patients are accrued to that treatment arm for a total of 25 patients per treatment arm.

**Definition of Evaluable Patients** 

Patients are considered evaluable for tumor response if they have completed at least 2 vaccinations and have undergone the tumor restaging at week 8.

Patients are considered evaluable for immune response if they have had at least 1 vaccination and have had immune analysis at week 4.

Patients are eligible for toxicity following a single vaccination.

#### Reporting of Outcomes

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Response rates are reported using descriptive statistics and report rates of CR, PR, SD, and PD in those patient determined to be evaluable. Time to progression following initial therapy is determined for those patients experiencing either a CR or PR.

Secondary endpoints include immune response, duration of response, and safety. These rates are also reported using descriptive statistics. Safety is reported as percent of patient experiencing a given adverse event. Mean time to progression for the responding population is reported using Kaplan-Meier statistics.

#### **Unmodified NSCLC Cell Lines**

Seven of the eight NSCLC lines used in the production of this vaccine are established cell lines that are purchased from American Tissue Cell Culture (ATCC). The human squamous NSCLC cell line, Rh-2, was established from a surgical resection specimen in the lab of Dr. Steven Dubinett at UCLA in 1994, and is publicly available per Lee et al., J. Immunology 152: 3222, 1994; Huang et al., Cancer Research 55: 3847, 1995; Huang et al., J. Immunology 157: 5512, 1996; and Huang et al., Cancer Research 58: 1208, 1998.

### pCHEK/HBA2:TGFβ2 Antisense Plasmid

The pCHEK vector used to construct the human TGF $\beta2$  antisense containing plasmid was derived from the pCEP4 vector (Invitrogen, San Diego, CA). It has been modified slightly to facilitate genetic subcloning. The resulting carrier vector is pCHEK. Genetic subcloning was used to insert the TGF $\beta2$  antisense gene fragment (HBA2) into pCHEK. Aliquots of the pCHEK/HBA2 plasmid were examined by restriction enzyme analyses to ensure 1) the identity of the carrier vector into which TGF $\beta2$  antisense was cloned and 2) the correct orientation of the TGF $\beta2$  antisense insert.

Test limits: Complete homology with expected DNA fragment sizes after a series of restriction digests with 14 endonucleases: Apal, BamHI, BgIII, Clal, EcoRV, HindIII, Hpal, Notl, Nrul, Pstl, Sall, Sacll, Scal, and Xbal.

Results: The observed DNA fragments obtained by these restriction digests corresponded with the expected fragment sizes of pCHEK/HBA2 plasmid. In conclusion, the vector used for subcloning is correct and the  $TGF\beta2$  antisense gene fragment insert is in the correct orientation.

# 30 The expected fragments of these restriction digests are:

Apal	$4604\; bp\;\; 3309\; bp$	1957 bp 874 bp	219 bp
BamHI	10963 bp		
BgIII	10210 bp	753 bp	
Clal	10963 bp		

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**EcoRV** 10963 bp HindIII 7540 bp 2787 bp 636 bp Hpal 10251 bp 712 bp Notl 10963 bp 5 Nrul 5716 bp 5247 bp Pstl 7573 bp 1494 bp 1277 bp 619 bp Sall 8738 bp 2225 bp SacII 5347 bp 3340 bp 2276 bp Scal 8021 bp 1915 bp 1027 bp 10 Xbal 9579 bp 1384 bp

#### TGF<sub>B</sub>2 Antisense Insert

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To further ensure the correct sequence and orientation of the TGF $\beta$ 2 antisense fragment, the insert was tested by sequence analyses using an ABI-310 Genetic Analyzer (Perkin Elmer, Foster City, CA).

Test limits: Complete homology between plasmid insert and TGF $\beta$ 2 antisense.

Results: Sequencing results obtained confirmed the presence of human TGF $\beta$ 2 fragment in the pCHEK vector. These results also confirmed the correct orientation of the insert. The sequence of the human TGF 2 fragment used in construction of the pCHEK/HBA2 vector and its flanking regions in the vector are as follows. Lower case letters represent the two vector sequences that flank human TGF $\beta$ 2 fragment.

20 tgtctggatc cggccttgcc ggcctcga (seq id no:2)-- vector sequence flanking the insert--

Base pair 5 of human TGFB2

# Gene-Modified NSCLC Cell Lines

Following transfection, clones of each gene-modified cell line were tested for the presence of pCHEK/HBA2 vector by PCR. Only those clones testing positive for pCHEK/HBA2 were chosen for further analysis.

Results: Colonies from seven of the eight cell lines tested were positive for TGF $\beta$ 2 antisense transfection. The cell line NCI-H-292 was negative.

#### TGFβ2 Downregulation

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Following transfection, clones of each gene-modified cell line were tested for TGF $\beta$ 2 downregulation by Enzyme Linked Immunosorbent Assay (ELISA) and compared to unmodified parental cell lines. Briefly, serum free supernatant of TGF $\beta$ 2 antisense gene modified cells cultures was collected after 24 hr and assayed in triplicate for TGF $\beta$ 2 secretion levels employing an ELISA kit (Genzyme, Cambridge, MASS). The human TGF $\beta$ 2 was captured by an anti-human TGF $\beta$ 2 monoclonal antibody and quantitated by reaction with horse radish peroxidase-conjugated goat anti-human TGF $\beta$ 2 antisera according to the manufacturer's recommendation. Quantitation was achieved by developing the enzymatic reaction with a chromagen substrate and reading the optical density on a micro-ELISA plate reader. A standard TGF $\beta$ 2 curve presenting known concentrations of TGF $\beta$ 2 permited quantitation of TGF $\beta$ 2 secretion by the TGF $\beta$ 2 antisense gene modified cells.

Test limits for unmodified tumor cells: Secretion of at least 200 pg TGF $\beta$ 2 /10<sup>6</sup> cells /24 hr.

Test limits for gene-modified tumor cell lines for vaccine: Lowering of TGF $\beta$ 2 production by at least 35% relative to unmodified parental cells. This test limit has been demonstrated to enhance immunogenicity of tumor cells in vaccination regimens, per Lee et al., J. Immunology 152: 3222, 1994.

Approximately 3-4 days prior to initiation of therapy, aliquots of each gene-modified cell line are re-tested for TGFβ2 inhibition. The same test limits apply.

Cell Line	Lung Carcinoma	Unmodified TGFB2  Levels  (ng/10° cell/24 hr)	Gene modified TGFB2  Levels  (ng/10° cell/24 hr)	% TGF \(\beta^2\) Down- regulation
NCI-H-292	Mucoepidermoid	ND	ND	ND
NCI-H-460	Large cell	0.67	0.12	82%
NCI-H-520	Squamous	2.27	1.43	37%
NCI-H-596	Adenosquamous	1.5	2.6	-73%
NCI-H-661	Large cell	13.1	12.96	1%
SK-LU-1	Adenocarcinoma	3.58	1.36	62%
SK-MES-1	Squamous	1.2	1.36	-13%
Rh-2	Squamous	1.16	0.1	91%

The 4 cell lines selected were: NCI-H-460, NCI-H-520, SK-LU-1 and Rh-2. In our hands, TGF $\beta$ 2 antisense gene modifications resulted in 37-91% blockage of intrinsic TGF $\beta$ 2 expression in tumor cells. Suppression of TGF $\beta$ 2 expression in these cells has been stable for period of six to nine months in culture.

# **Cell Density**

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Prior to patient vaccination cell density is assessed for each cell line. An aliquot of each cell line to be used is counted using a hemocytometer to ascertain cell density. Equal numbers of each gene-modified cell line is admixed for the three doses to be investigated in this study  $(1.25 \times 10^6, 2.5 \times 10^6 \text{ and } 5 \times 10^6 \text{ total cells respectively})$ .

#### **Cell Viability**

Prior to patient vaccination cell viability is also assessed for each cell line. The viability of cells employed for immunization is evaluated by trypan blue exclusion methods. Trypan blue dye is a measure of plasma membrane integrity. Viable cells maintain plasma membrane integrity and therefore exclude the dye. Dead cells lose membrane integrity and allow uptake of the dye thus appearing blue. An aliquot of each cell line is tested and cells counted in a hemocytometer. The percentage of viable "non-blue" cells is determined.

Test Limits: The viability of the TGF $\beta$ 2 antisense gene modified tumor cells used for therapy must be greater than 50%.

### Clonogenicity

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To ensure safety, all gene-modified tumor cell lines to be used in patient vaccinations must be irradiated prior to injection. This is to prevent tumor cell growth and replication. Cells are irradiated prior to use with a dose of 10,000 cGy. The selection of this radiation dose is based on discussions with Dr. Herman Suit, Chief of Radiation Oncology at Massachusetts General Hospital. This was the lowest radiation dose sufficient to render the tumor cells incapable of proliferation and tumor formation. It is our desire to utilize the lowest possible radiation dose for the transfected cells to optimize the level and duration of TGFβ2 antisense transcription. In addition, we have tested this irradiation dose in our laboratory on cultured tumor cells of different histologic origins, including human NSCLC, gliomas, colon cancer, and pancreatic carcinoma cell lines and demonstrated that it is capable of completely arresting colony formation by cultured tumor cells of different histologic origins.

Samples of unmodified and gene modified human NSCLC cell lines were irradiated with 10,000 centi-Grays. The irradiated cells were then cultured in T-225 flasks and observed for colony formation. A colony was defined as a cluster of 16 growing cells. As presented in the table below, colonies did not form in the irradiated cultures during a four-six week observation period. In contrast, all the non-irradiated control cultures became confluent after 10-14 days. Cell death occurred approximately two weeks after initiation of the irradiated cultures.

Effect of radiation on primary NSCLC cell cultures

Tumor cells	Radiation Dose (Gys)	# Colonies at 5 weeks
Control cultures	None	Confluent after 10-14 days
NCI-H-292	10,000	None
NCI-H-460	10,000	None
NCI-H-520	10,000	None
NCI-H-596	10,000	None
NCI-H-661	10,000	None
SK-Lu 1	10,000	None
SK-MES-1	10,000	None
Rh-2	10,000	None

Prior to vaccination, an aliquot of each gene-modified cell line is thawed and tested for colony formation during a four to six week culturing period before each lot is deemed safe for patient injection.

Test Limits: No colony formation.

#### Sterility of Cell Lines

Sterility testing was performed for each unmodified cell line by ATCC, the manufacturer of the cells.

In addition, aliquots of each line were sent to Molecular Diagnostics Associates to assay for the presence of the following viral agents:

5 HIV 1 & 2 HBV CMV
HH-6 HCV HTLV
EBV Adventitious viruses

10 Results: All eight unmodified master cell lines were found to be negative for the presence of bacteria, fungi and viruses.

During *in vitro* growth and manipulations each cell line was routinely tested for bacterial, fungal, and mycoplasma infection. To avoid contamination with other cells, cultures were processed individually at all points during laboratory manipulations: Finally, on the day of therapy, a sample of the inoculum is retested by a gram stain. Only cells that pass all sterility testing are used for therapy.

Test limits are: No bacteria, mycoplasma or fungal infections.

#### Clinical Grade pCHEK/HBA2 Plasmid

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Following preliminary characterization of pCHEK/HBA2 plasmid, DNA stocks were prepared from bacterial cultures by the alkaline lysis method of Birnboim and Doly as optimized by Qiagen Corporation (Birnboim and Doly, 1979), and purified on Qiagen EndoFree Giga prep columns. All steps were carried out under sterile conditions using ART barrier tips in the biosafety flow hood. An aliquot of the plasmid DNA was removed and tested for sterility. Briefly 20  $\mu$ l of plasmid DNA was used to inoculate four culture tubes each containing 5 ml of antibiotic free LB. The cultures were incubated for five days at 37 °C.

Test Limits: No bacteria growth.

Results: No colonies were observed confirming the sterility of the prepared clinical grade DNA.

# **Brief General Description of Manufacturing and Packaging Procedures**

Eight established NSCLC cell lines were purchased from American Tissue Cell Culture (ATCC) or otherwise and were expanded and frozen as unmodified Master Cell Banks (unMCB). Each line was tested for sterility (bacterial and viral contaminants), clonogenicity and TGFβ2 expression. Aliquots of each line were thawed and transfected with pCHEK/HBA2, a vector containing the TGFβ2 antisense transgene, using standard techniques. Gene-modified cell lines were expanded in culture, under hygromycin selection, to grow sufficient numbers for therapeutic applications and testing. Expanded cell lines were then assayed for down-regulation of TGFβ2 expression and sterility. Four NSCLC cell lines which demonstrated successful downregulation of TGFβ2 expression and sterility were identified: NCI-H-460, NCI-H-520, SK-LU-1 and Rh-2. These cell lines were frozen in aliquots as (1) gene-modified Master Cell Banks (gmMCB) and (2) gene-modified Working Cell Banks (gmWCB) and stored in liquid nitrogen. Before use, aliquots of

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these four cell lines are thawed from the gmWCB, irradiated with 10,000 Gy and re-tested for sterility, clonogeneicity and  $TGF\beta2$  downregulation. Only cell lots passing all test limits are acceptable for vaccine preparation. On the day of injection, sufficient cells from each of the acceptable gmWCB lots are then thawed, irradiated and admixed in equal numbers. Before patient vaccination, a sample of the innoculum is tested for bacterial contamination. If no contamination is detected, vaccination can proceed.

# **Tissue Procurement**

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The following eight established NSCLC cell lines were obtained from American Type Culture Collection (ATCC) or otherwise, expanded in culture and frozen as the unmodified Master Cell Banks (Total 8 unMCB). Cell lines were cultured in IMDM supplemented with 10% FBS, 25 mM Hepes, 2 mM L-glutamine, 1 mM sodium pyruvate, 2.5  $\mu$ g/ml fungizone, 50  $\mu$ g/ml gentamycin sulphate, 10<sup>4</sup> M  $\alpha$ -thio-glycerol and non-essential amino acids. Each cell line was frozen as one lot containing 100 vials at  $> 10^8$  cells/vial. Each line was tested for sterility and TGF 2 expression. The following cell lines were used:

- NCI-H-292
- NCI-H-460
- 15 NCI-H-520
  - NCI-H-596
  - NCI-H-661
  - SK-LU-1
  - SK-MES-1
- 20 Rh-2

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#### Construction of Human TGFB2 Antisense Expression Plasmid

### PCHEK/HBA2 Plasmid Description

The pCHEK vector used to construct the human TGF $\beta$ 2 antisense expression plasmid (pCHEK/HBA2) was derived from the pCEP4 vector (Invitrogen, San Diego, CA) to facilitate gene modification of cancer cells and to eliminate safety concerns. The pCHEK vector is identical to the pCEP4 vector in all regions except the following:

- Kanamycin resistance, instead of the ampicillin resistance is incorporated into the pCHEK vector.
- In the pCHEK vector, a DNA cassette unit consisting of the SV-40 early promoter followed by an intron
  drives the expression of the hygromycin resistance gene. Incorporation of the SV-40 promoter/intron unit is
  to increase expression of the hygromycin resistance gene to facilitate selection of gene modified cells in
  culture.

The pCHEK/HBA2 plasmid utilizes a CMV promoter to drive the expression of a 930 base pair human TGF $\beta$ 2 fragment in antisense orientation. The TGF $\beta$ 2 antisense fragment consists of bases 6-935 of the 5' end of the human

TGFβ2 cDNA molecule that was ligated in reverse orientation adjacent to and under the control of the CMV promoter. The pCHEK vector also contains the hygromycin resistance gene driven by the SV-40 early promoter, the Epstein-Barr virus origin of replication, and the gene for the Epstein-Barr virus nuclear associated protein 1 (EBNA-1). Additionally, the vector contains the ColE1 origin and kanamycin resistance genes for selection of bacteria containing vector during DNA manufacture.

#### **PCHEK VECTOR DOMAINS**

Domain	Fragment containing the domain
SV-40 Poly A signal	1-405
Multiple cloning site	406-463
CMV promoter	467-1311
TK poly A signal	1312-1843
Hygromycin	1844-2899
Intron	2900-3233
SV-40 early promoter	3234-3602
ColE1 Origin & Kanamycin resistance	3603-5188
EBNA-1	5189-7789
Epstein Barr origin of replication (OriP)	7790-1046

Subcloning TGF\(\beta\)2 Antisense into pCHEK Vector

TGFβ2 Isolation

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To construct the pCHEK/HBA2 plasmid we first constructed pCEP4/HBA2, a shuttle vector. Briefly, the plasmid pPC21 (publicly available from Dr. Purchio) containing the human TGF $\beta$ 2 gene was digested to completion with EcoRI. The EcoRI ends were blunt ended by adjusting the reaction conditions to 250  $\mu$ M each dATP, dCTP, dGTP, and dTTP and adding 3 units Klenow enzyme to initiate the reaction. The reaction was allowed to proceed for 30 minutes at 37 °C. The reaction volume was then adjusted to 100  $\mu$ l with TE and phenol/chloroform extracted. The DNA was isopropanol precipitated and rinsed with 70% ethanol. The 930 base pair TGF $\beta$ 2 fragment (HBA2) was released from the vector by Hind III digestion. Following electrophoresis in a 1% agarose gel, a gel slice containing the 930 bp TGF $\beta$ 2 fragment was excised and the DNA fragment extracted by routine oxidized silica (glass powder) method. The TGF $\beta$ 2 DNA was then ready to be ligated into the pCEP4 vector.

#### pCEP4/HBA2 Shuttle Vector Construction

The pCEP4 vector was prepared by digestion with Xhol restriction enzyme, and blunt ended by Klenow reaction as described above. Following phenol/chloroform extraction and ethanol precipitation, the vector was digested with Hind III and purified by agarose gel electrophoresis / glass powder method as described above.

The 930 base pair human TGFβ2 fragment was directionally subcloned into the pCEP4 vector in antisense orientation. Following transformation of E. coli, pCEP4/HBA2 DNA was prepared from several ampicillin resistant transformed bacterial colonies. The isolated DNA from these colonies was characterized by restriction enzyme analysis and one clone designated as pCEP4/HBA2 was selected and used for construction of the clinical plasmid pCHEK/HBA2.

pCHEK/HBA2 Expression Plasmid Construction

Briefly, the pCEP4/HBA2 DNA was digested with restriction enzymes Kpn I and Bam HI and the 930 bp TGF 2 antisense insert fragment was purified by agarose gel electrophoresis. The insert was then ligated into the Kpn I and Bam HI digested pCHEK vector, and used to transform bacteria. Following kanamycin selection of overnight culture, pCHEK/HBA2 DNA was isolated from several clones and characterized by restriction enzyme analyses to ensure correct identity. To further ensure the correct sequence and orientation of the TGFβ2 antisense fragment, the insert was tested by sequence analyses using an ABI-310 Genetic Analyzer (Perkin Elmer, Foster City, CA).

#### Manufacturing of Clinical Grade pCHEK/HBA2 Plasmid

Following preliminary characterization, one bacterial colony containing the pCHEK/HBA2 plasmid was streaked on a Luria-Bertani (LB) agar plate containing 100  $\mu$ g/ml kanamycin. Following incubation at 37 °C, a single bacterial colony was used to inoculate 5 ml of LB broth containing 100  $\mu$ g /ml kanamycin and grown overnight at 37 °C in a shaking bacterial incubator. This overnight culture was used to inoculate 50 ml culture of plasmid containing bacteria. The 50 ml bacterial culture was incubated overnight and used to inoculate flasks containing 2 liters of LB plus 100  $\mu$ g /ml kanamycin. This was grown overnight at 37 °C.

DNA was prepared from bacterial cultures by the alkaline lysis method of Birnboim and Doly as optimized by Qiagen Corporation (Birnboim and Doly, 1979), and purified on Qiagen EndoFree Giga prep columns. All steps were carried out under sterile conditions using ART barrier tips in the biosafety flow hood.

An aliquot of DNA was removed for restriction analysis and determination of DNA concentration. The plasmid DNA concentration was adjusted to one mg/ml, divided into aliquots, and stored at -70 °C for future use. An aliquot of the plasmid DNA was removed and tested for sterility.

# Genetic Modification of NSCLC Tumor Cell Lines with pCHEK/HBA2 Plasmid

Four aliquots of each NSCLC cell line were removed from the appropriate unMCB and grown in culture. Cells were fed with fresh medium twice a week. On the day of gene modification cells were fed with fresh medium. Four hours later cells were trypsinized, washed with serum containing medium and subsequently PBS. Cell density was adjusted to  $1.2 \times 10^7$  cells per ml in a volume of 350  $\mu$ l PBS and incubated on ice for 15-20 minutes. 50  $\mu$ g pCHEK/HBA2 plasmid was added. The mixture was incubated on ice for an additional 10-15 minutes. The cell suspension was then transferred to a pre-chilled cuvette and incubated on ice. After five minutes the cuvettes were

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placed in a square wave electroporator (Genetronics, San Diego, CA) and subjected to three electroporation pulses of 3000 v/cm each pulse lasting 75 µseconds. 1 ml of cold fresh media contining 50 mM Hepes was added and the mixture incubated at room temperature for 10 minutes. Cells were then plated for 2 division cycles and then selection was begun (72 hours after transfection). Fresh medium containing 25 µg hygromycin/ml was added to the cultures. The gene-modified cells were expanded in culture and frozen as gene modified Master Cell Banks (gmMCB) for a total of 4 gmMCBs per cell line. The presence of pCHEK/HBA2 plasmid in gene modified cells was ascertained by PCR. In addition, five vials (5%) from each gmMCB were submitted and used for sterility testing consisting of aerobic, anaerobic, mycoplasma, and fugal assays.

#### Preparation and Identification of Cell Lines for Patient Vaccination

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Following genetic modification, tumor cell lines were expanded in culture to grow sufficient cells for therapeutic applications and testing. Clones from each cell line passing identity, strength and safety test limits were cryopreserved in liquid nitrogen as aliquots of gene-modified Master Cell Banks (gmMCB) and gene-modified Working Cell Banks (gmWCB). Prior to patient vaccination, aliquots from each gmWCB lot are thawed and irradiated with a dose of 10,000 cGy, a radiation dose we have demonstrated to be capable of completely arresting colony formation by cultured tumor cells of different histologic origins including these NSCLC cells. Aliquots from each lot undergo safety, strength and identity tests to ensure alteration or contamination has not occurred during cell manipulations and freezing. Lots are tested for: clonogenicity, sterility and TGFβ2 downregulation.

# Cell Preparation and Testing Prior to Patient Vaccination

Prior to subcutaneous immunization, aliquots of the four chosen cell lines are placed in short term cultures. Gene-modified cells are detached from culture dishes, washed and resuspended in medium, irradiated with 10,000 cGy, washed and resuspended in lactated Ringer's solution. They are then tested for sterility and viability. Only cells passing test limits are used for patient vaccination.

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While the present invention has been described in some detail for purposes of clarity and understanding, one skilled in the art will appreciate that various changes in form and detail can be made without departing from the true scope of the invention. All figures, tables, and appendices, as well as patents, applications, and publications, referred to above, are hereby incorporated by reference.

#### WHAT IS CLAIMED IS:

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1. A composition comprising a therapeutically effective amount of genetically modified cells containing a genetic construct expressing a TGF $\beta$  inhibitor effective to reduce expression of TGF $\beta$ , wherein said genetically modified cells are lung cancer cells.

- 2. Use of a composition comprising a therapeutically effective amount of genetically modified cells containing a genetic construct expressing a TGF $\beta$  inhibitor effective to reduce expression of TGF $\beta$ , wherein said genetically modified cells are lung cancer cells for the preparation of a medicament for prolonging survival of a subject having a lung cancer.
- 3. A method for prolonging survival of a subject having a lung cancer comprising the step of administering to said subject a composition comprising a therapeutically effective amount of genetically modified cells containing a genetic construct expressing a TGF $\beta$  inhibitor effective to reduce expression of TGF $\beta$ , wherein said genetically modified cells are lung cancer cells.
- 4. The composition, use, or method of any of Claims 1-3, wherein said lung cancer cells are non-small cell lung cancer (NSCLC) cells.
- 15 5. The composition, use, or method of any of Claims 1-3, wherein said lung cancer cells are small cell lung cancer (SCLC) cells.
  - 6. The composition, use, or method of any of Claims 1-3, wherein said TGF $\beta$  is TGF $\beta$ -1.
  - 7. The composition, use, or method of any of Claims 1-3, wherein said TGF $\beta$  is TGF $\beta$ -2.
  - 8. The composition, use, or method of any of Claims 1-3, wherein said TGF $\beta$  is TGF $\beta$ -3.
- 20 9. The composition, use, or method of any of Claims 1-3, wherein said genetically modified cells are autologous cells.
  - 10. The composition, use, or method of any of Claims 1-3, wherein said genetically modified cells are allogeneic cells.
  - 11. The composition, use, or method of any of Claims 1-3, wherein said genetically modified cells are mixtures of autologous and allogeneic cells.
  - 12. The composition, use, or method of any of Claims 1-3, wherein said genetically modified cells further express one or more cytokines having immunostimulatory effects.
  - 13. The composition, use, or method of Claim 12, wherein said one or more cytokines is selected from the group consisting of interleukin-1, interleukin-2, interleukin-3, interleukin-4, interleukin-5, interleukin-6, interleukin-7, interleukin-8, interleukin-9, interleukin-10, interleukin-11, interleukin-12, interleukin-15, interferon-alpha, interferongamma, tumor necrosis factor-alpha, transforming growth factor-beta, granulocyte macrophage colony stimulating factor, and granulocyte colony stimulating factor.
    - 14. The composition, use, or method of any of Claims 1-3, wherein said  $TGF\beta$  inhibitor is an antisense.

The composition, use, or method of any of Claims 1-3, wherein said TGF $\beta$  inhibitor is an antisense having the sequence of SEQ ID NO:1.

- 16. The composition, use, or method of any of Claims 1-3, wherein said  $TGF\beta$  inhibitor is a ribozyme.
- The composition, use, or method of any of Claims 1-3, wherein said TGF $\beta$  inhibitor is a dominant negative mutant.
  - 18. The composition, use, or method of any of Claims 1-3, wherein said TGF $\beta$  inhibitor is an antibody.
  - 19. The composition, use, or method of any of Claims 1-3, wherein said  $TGF\beta$  inhibitor prevents  $TGF\beta$  receptor and Smad protein interaction.
- The composition, use, or method of any of Claims 1-3, wherein said composition is provided in unit dosage form.

#### SEQUENCE LISTING

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